

L6 ANSWER 24 OF 34 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on
STN
AN 1993:428737 BIOSIS
DN PREV199396083362
TI Engineering proteins for nonnatural environments.
AU Arnold, Frances H.
CS Div. Chem. Chemical Eng., Calif. Inst. Technol., Pasadena, CA 91125, USA
SO FASEB (Federation of American Societies for Experimental Biology) Journal,
(1993) Vol. 7, No. 9, pp. 744-749.
CODEN: FAJOEC. ISSN: 0892-6638.
DT Article
LA English
ED Entered STN: 22 Sep 1993
Last Updated on STN: 22 Sep 1993
AB The ability to use proteins in nonnatural environments greatly expands their potential applications in biotechnology. Because nature has not paid much attention to optimizing proteins for *in vitro* applications under conditions that differ substantially from their natural surroundings, there is generally room for improvement through alterations in the amino acid sequence. The most effective approach to this protein engineering task depends on the level to which the molecular basis for the desired property is understood. Consistently successful "rational" design using site-directed mutagenesis requires a high level of understanding of structure and mechanisms or, alternatively, a particularly simple strategy for obtaining the desired feature. An example of a generally applicable and easy-to-implement protein stabilization strategy is metal ion chelation by specific surface dihistidine sites, which can affect thermal stability as well as the protein's ability to withstand denaturants such as guanidinium chloride. Random mutagenesis, on the other hand, can be effective even when structure or mechanisms are poorly understood, provided one can conveniently screen or select for the property of interest. This approach is illustrated by the sequential accumulation of random mutations that greatly enhance the catalytic activity of a serine protease, subtilisin E, in polar organic solvents. The random mutagenesis approach, which mimics the natural evolutionary refinement process, can be used to "coax" enzymes into tolerating nonnatural environments.

L11 ANSWER 48 OF 49 MEDLINE on STN

DUPLICATE 22

AN 82280203 MEDLINE

DN PubMed ID: 7051707

TI [Thermitase, a thermostable **serine protease** of Thermoactinomyces vulgaris: interaction of the active center and the SH-group of the enzyme].

Thermitase, eine thermostabile Serin-Protease aus Thermo- actinomyces vulgaris: Wechselwirkung zwischen aktivem Zentrum und SH-Gruppe des Enzyms.

AU Hansen G; Frommel C; Hausdorf G; Bauer S

SO Acta biologica et medica Germanica, (1982) 41 (2-3) 137-44.

Journal code: 0370276. ISSN: 0001-5318.

CY GERMANY, EAST: German Democratic Republic

DT Journal; Article; (JOURNAL ARTICLE)

LA German

FS Priority Journals

EM 198210

ED Entered STN: 19900317

Last Updated on STN: 20000303

Entered Medline: 19821021

AB Modification of the serine and histidine residue in the active centre of thermitase with diisopropylfluorophosphate (DFP) or L-1-tosylamide-2-phenylethyl chloromethylketon (TPCK), and of the only SH-group of the enzyme, with Hg-compounds causes an activity loss against hydrolysis of 4-nitrophenylacetate. While the **modification** of **cysteine** prevents reaction of **serine** and histidine in the active centre of the enzyme with DFP and TPCK, respectively, the Hg²⁺- and CF₃Hg⁺-binding to the SH-group after modification of essential amino acid residues in the active centre is retained. To elucidate the interaction of the SH-group with the active centre, the modified products of thermitase were investigated for their thermostability. Ca²⁺-ions were found to have a stabilizing effect on all the modified products of thermitase, as well as on the native enzyme. Simultaneous **modification** of the **cysteine** and **serine** leads to an increase in thermostability of thermitase, whilst double **modification** at the cysteine and histidine causes destabilization of the enzyme.

Database : Issued_Patents_AA:*

1: /cgn2_6/ptodata/2/iaa/5A_COMB.pep:*

2: /cgn2_6/ptodata/2/iaa/5B_COMB.pep:*

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5: /cgn2_6/ptodata/2/iaa/PCTUS_COMB.pep:*

6: /cgn2_6/ptodata/2/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
No.	Score	Match	Length	DB	ID	%
1	4681	100.0	855	2	US-09-027-337-2	5972616
2	4681	100.0	855	4	US-09-644-600-2	6451500
3	4681	100.0	855	4	US-09-654-600A-2	6649741
4	3810	81.4	902	4	US-09-644-600-10	
5	3810	81.4	902	4	US-09-654-600A-10	
6	703.5	15.0	798	1	US-08-200-900A-2	
7	703.5	15.0	798	5	PCT-US94-00616-2	
8	588	12.6	407	4	US-09-734-675-4	
9	560.5	12.0	492	4	US-09-685-166A-895	
10	558.5	11.9	492	3	US-09-342-749-2	
11	558.5	11.9	492	4	US-09-691-840-2	
12	547.5	11.7	235	3	US-08-944-483-65	
13	544.5	11.6	235	3	US-08-807-151-3	
14	544.5	11.6	235	4	US-09-478-957-3	

Db	421	NSNKITVRFHSDQSYTDTGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH	480
Qy	481	SDELNCSCDAGHQFTCKNFKPLFWVCDSVNDCGDSDEQGCSCPAQTFRCSNGKCLSK	540
Db	481	SDELNCSCDAGHQFTCKNFKPLFWVCDSVNDCGDSDEQGCSCPAQTFRCSNGKCLSK	540
Qy	541	SQQCNGKDDCGDSDEASCPKVNVTCTKHTYRCLNGLCLSKGPNPECDGKEDCSDGSDEK	600
Db	541	SQQCNGKDDCGDSDEASCPKVNVTCTKHTYRCLNGLCLSKGPNPECDGKEDCSDGSDEK	600
Qy	601	DCDCGLRSFTRQARVVGTDADEGEWPWQVSLHALGQGHICASLISPWNWLVSAAHCYID	660
Db	601	DCDCGLRSFTRQARVVGTDADEGEWPWQVSLHALGQGHICASLISPWNWLVSAAHCYID	660
Qy	661	DRGFYRSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP	720
Db	661	DRGFYRSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP	720
Qy	721	AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCEENLL	780
Db	721	AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCEENLL	780
Qy	781	PQQITPRMMCVGFLSGGVDSQGDSSGGLSSVEADGRIFQAGVVSWGDGCAQRNKGVYT	840
Db	781	PQQITPRMMCVGFLSGGVDSQGDSSGGLSSVEADGRIFQAGVVSWGDGCAQRNKGVYT	840
Qy	841	RLPLFRDWIKENTGV	855
Db	841	RLPLFRDWIKENTGV	855

RESULT 2

US-09-644-600-2

; Sequence 2, Application US/09644600

; Patent No. 6451500

; GENERAL INFORMATION:

; APPLICANT: O'Brien, Timothy J.

; APPLICANT: Tanimoto, Hirotoshi

; TITLE OF INVENTION: TADG-15: An Extracellular Serine Protease

; TITLE OF INVENTION: Overexpressed in Carcinomas

; FILE REFERENCE: D6064CIP/D

; CURRENT APPLICATION NUMBER: US/09/644,600

; CURRENT FILING DATE: 2000-08-23

; PRIOR APPLICATION NUMBER: 09/421,213

; PRIOR FILING DATE: 1999-10-20

; PRIOR APPLICATION NUMBER: 09/027,337

; PRIOR FILING DATE: 1998-02-20

; NUMBER OF SEQ ID NOS: 98

; SEQ ID NO 2

; LENGTH: 855

; TYPE: PRT

; ORGANISM: Homo sapiens

; FEATURE:

; OTHER INFORMATION: TADG-15

US-09-644-600-2

Query Match 100.0%; Score 4681; DB 4; Length 855;
 Best Local Similarity 100.0%; Pred. No. 0;
 Matches 855; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy	1	MGSDRARKGGGGPKDFGAGLKYNRSRHEKVNGLEEGVEFLPVNNVKVEKHGPGRWVVLAA	60
Db	1	MGSDRARKGGGGPKDFGAGLKYNRSRHEKVNGLEEGVEFLPVNNVKVEKHGPGRWVVLAA	60
Qy	61	VLIGLLLVLIGFLVWHLQYRDVRVQKVFNNGYMRITNENFDAYENSNSTEFVSLASKV	120
Db	61	VLIGLLLVLIGFLVWHLQYRDVRVQKVFNNGYMRITNENFDAYENSNSTEFVSLASKV	120
Qy	121	KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM	180
Db	121	KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM	180
Qy	181	LPPRARSLKSFVVTSVVAFPTDSKTVQRTQDNCSFGLHARGVELMRFTTPGFPDSPYPA	240
Db	181	LPPRARSLKSFVVTSVVAFPTDSKTVQRTQDNCSFGLHARGVELMRFTTPGFPDSPYPA	240
Qy	241	HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYNTLSPMEPHALVQLCGTYPPS	300
Db	241	HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYNTLSPMEPHALVQLCGTYPPS	300
Qy	301	YNLTFHSSQNVLLITLITNTERRHPGFEATFFQLPRMSSCGGRLRKAQGTFNSPYYPGHY	360
Db	301	YNLTFHSSQNVLLITLITNTERRHPGFEATFFQLPRMSSCGGRLRKAQGTFNSPYYPGHY	360
Qy	361	PPNIDCTWNIEVPNNQHVVKVSKFFYLLEPGVPAGTCPKDVEINGEKYCGERSQFVVTS	420
Db	361	PPNIDCTWNIEVPNNQHVVKVSKFFYLLEPGVPAGTCPKDVEINGEKYCGERSQFVVTS	420
Qy	421	NSNKITVRFHSDQSYTDGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH	480
Db	421	NSNKITVRFHSDQSYTDGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH	480
Qy	481	SDELNCSCDAGHQFTCKNFKPLFWVCDSVNDGDNDEQGCSCPAQTFRCNSNGKCLSK	540
Db	481	SDELNCSCDAGHQFTCKNFKPLFWVCDSVNDGDNDEQGCSCPAQTFRCNSNGKCLSK	540
Qy	541	SQQCNGKDDCGDGSDEASCPKVNVVTCTKHTYRCLNGLCLSKGNECDGKEDCSDGSDEK	600
Db	541	SQQCNGKDDCGDGSDEASCPKVNVVTCTKHTYRCLNGLCLSKGNECDGKEDCSDGSDEK	600
Qy	601	DCDCGLRSFTRQARVVGGTDADEGEWPWQVSLHALGQGHICGASLISPWLVSAAHYID	660
Db	601	DCDCGLRSFTRQARVVGGTDADEGEWPWQVSLHALGQGHICGASLISPWLVSAAHYID	660
Qy	661	DRGFRYSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP	720
Db	661	DRGFRYSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP	720
Qy	721	AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCENLL	780
Db	721	AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCENLL	780
Qy	781	PQQITPRMMCVGFLSGGVDSQGDGGPLSSVEADGRIFQAGVVSWGDGCAQRNKPGVYT	840

Db 781 ||||||| PQQITPRMMCVGFLSGGVVDSCQGDSGGPLSSVEADGRIFQAGVVSWGDGCAQRNKGPGVYT 840
Qy 841 RLPLFRDWIKENTGV 855
Db 841 RLPLFRDWIKENTGV 855

RESULT 3

US-09-654-600A-2

; Sequence 2, Application US/09654600A
; Patent No. 6649741
; GENERAL INFORMATION:
; APPLICANT: O'Brien, Timothy J.
; APPLICANT: Tanimoto, Hirotoshi
; TITLE OF INVENTION: TADG-15: An Extracellular Serine Protease
; TITLE OF INVENTION: Overexpressed in Carcinomas
; FILE REFERENCE: D6064CIP/D
; CURRENT APPLICATION NUMBER: US/09/654,600A
; CURRENT FILING DATE: 2000-09-01
; PRIOR APPLICATION NUMBER: 09/421,213
; 09/027,337
; PRIOR FILING DATE: 1999-10-20
; 1998-02-20
; NUMBER OF SEQ ID NOS: 98
; SEQ ID NO 2
; LENGTH: 855
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: TADG-15

US-09-654-600A-2

Query Match 100.0%; Score 4681; DB 4; Length 855;
Best Local Similarity 100.0%; Pred. No. 0;
Matches 855; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 MGSDRARKGGGGPKDFGAGLKYNNSRHEKVNGLEEGVEFLPVNNVKVEKGPGRWVLAA 60
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Qy 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM 180
Db 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM 180

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Qy 421 NSNKITVRFHSDQSYTDTGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH 480
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Db 661 DRGFRYSDPTQWTAFLGLHDQSQR SAPGVQERRLKRIISHPFFNDFTFDYDIALLELEKP 720

Qy 721 AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTT CENLL 780
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Qy 781 PQQITPRMMC VGF LSGGV DSCQGD SGGPLSSVEADGRIFQAGV VSWGDGCAQRN KPGVYT 840
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Qy 841 RLPLFRDWIKENTGV 855
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Db 841 RLPLFRDWIKENTGV 855

RESULT 1

JC7731

membrane-bound arginine-specific serine proteinase precursor - rat

C;Species: Rattus norvegicus (Norway rat)

C;Date: 14-Dec-2001 #sequence_revision 14-Dec-2001 #text_change 03-Feb-2003

C;Accession: JC7731; JC7775

R;Kishi, K.; Yamazaki, K.; Yasuda, I.; Yahagi, N.; Ichinose, M.; Tsuchiya, Y.; Athauda, S.B.P.; Inoue, H.; Takahashi, K.

J. Biochem. 130, 425-430, 2001

A;Title: Characterization of a membrane-bound arginine-specific serine protease from rat intestinal mucosa.

A;Reference number: JC7731; MUID:21421307; PMID:11530019

A;Accession: JC7731

A;Molecule type: mRNA

A;Residues: 1-855 <KIS>

A;Cross-references: DDBJ:AB049189

A;Experimental source: strain Male, 7-week-old

R;Satomi, S.; Yamasaki, Y.; Tsuzuki, S.; Hitomi, Y.; Iwanaga, T.; Fushiki, T. Biochem. Biophys. Res. Commun. 287, 995-1002, 2001

A;Title: A role for membrane-type serine protease (MT-SP1) in intestinal epithelial turnover.

A;Reference number: JC7775; PMID:11573963

A;Contents: Small intestine

A;Accession: JC7775

A;Molecule type: mRNA

A;Residues: 1-855 <SAT>

A;Cross-references: DDBJ:AB037898

C;Comment: This enzyme, an epithelial-derived, type II integral membrane serine protease. It localized mainly on brushborder membranes of the intestine and participates in the processing or digestion of specific proteins or peptides on the brushborder membranes. It also participates in the control of intestinal epithelial turnover by regulating the cell-substratum adhesion associated with epithelial migration and/or cell loss.

C;Genetics:

A;Gene: mt-spl

A;Map position: basolateral cell surface

C;Superfamily: membrane-bound arginine-specific serine proteinase

C;Keywords: protein digestion

Query Match 83.0%; Score 3883; DB 2; Length 855;
Best Local Similarity 81.1%; Pred. No. 4.5e-247;
Matches 693; Conservative 79; Mismatches 83; Indels 0; Gaps 0;

Qy 1 MGSDRARKGGGGPKDFGAGLKYNRSRHEKVNGLEEGVEFLPVNNVKVKEHGPGRWVVLAA 60
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Db 61 VVFSFLLSLMAGLLVWHFHYRNVRIQKVFNGHLRITNENFLDAYENSTSTEFISLASQV 120

Qy 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM 180
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Db 121 KEALKLMLYSEVPVLGPYHKSTVTAFSEGSVIAYWSEFSIPPHLEEEVDRAMAVERVVT 180

Qy 181 LPPRARSLKSFVVTSVVAFPTDSKTVQRTQDNCSFGLHARGVELMRFTTPGFPDSPYPA 240
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Db 181 LPPRARALKSFVLTSVVAFPIDPRMLQRTQDNSCSFALHARGRTVTRFTPGFPNSPYPA 240

Qy 241 HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYNTLSPMEPHALVQLCGTYPPS 300
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Db 241 HARCQWVLRGDADSVLSLTFRSFDVAPCDGHDSLVTYDSLSPMEPHAVVRLCGTFSPS 300

Qy 301 YNLTFHSSQNVLLITLITNTERRHPGEATFFQLPRMSSCGGRLRKAQGTFNSPYYPGHY 360
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Db 301 YNLTFLSSQNVFLVTLITNTDRRHPGEATFFQLPKMSSCGLLSEAQGTFSSPYYPGHY 360

Qy 361 PPNIDCTWNIEVPNNQHVVKVSFKFFYLLEPGVPAGTCPKDYVEINGEKYCGERSQFVVTS 420
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Db 361 PPNINCTWNVIKVPNNRNVKVRFKLFYLVDPNIPVGSCTKDYVEINGEKFCGERSQFVVSS 420

Qy 421 NSNKITVRFHSDQSYTDGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH 480
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Db 421 NSSKITVHFHSDHSYTDGFLAEYLSYDSNDPCPGFMCKTGRCIRKDLRCDGWADCPDY 480

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Qy 541 SQQCNGKDDCGDSDEASCPKVNVVTCTKHTYRCLNGLCLS KGNPECDGKEDCSDGSDEK 600
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Qy 601 DCDCGLRSFTRQARVVGGTDADEGEWPWQVSLHALGQGHICGASLISPNWLVSAAHCYID 660
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Db 601 NCDCGLRSFTKQARVVGGTNADEGEWPWQVSLHALGQGHLCGASLISPDWLVSAAHCFQD 660

Qy 661 DRGFRYSDPTQWTAFLGLHDQSQRSAPGVQERRLKRIISHPFFNDFTFDYDIALLELEKP 720
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Db 661 ETIFKYSDDHTMWTAFGLLDDQSKRSASGVQEHLKRIIITHPSFNDFTFDYDIALLELEKP 720

Qy 721 AEYSSMVRPICLPDASHVFpagkaiWVTGWGHTQYGGTGALILQKGEIRVINQTTCEENLL 780
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Db 721 AEYSTVVRPICLPDNTHVFpagkaiWVTGWGHTKEGGTGALILQKGEIRVINQTTCEELL 780

Qy 781 PQQITPRMMCVGFLSGGVVDSCQGDGGPLSSVEADGRIFQAGVVSWGDGCAQRNKPGVYT 840
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Db 781 PQQITPRMMCVGFLSGGVVDSCQGDGGPLSSVEKDGRIFQAGVVSWGECAQRNKPGVYT 840

Qy 841 RLPLFRDWIKENTGV 855
| :| | ||| | | |

Db 841 RIPEVRDWIKEQTGV 855

Database : SPTREMBL_25:*

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2: sp_bacteria:*

3: sp_fungi:*

4: sp_human:*

5: sp_invertebrate:*

6: sp_mammal:*

7: sp_mhc:*

8: sp_organelle:*

9: sp_phage:*

10: sp_plant:*

11: sp_rat:*

12: sp_virus:*

13: sp_vertebrate:*

14: sp_unclassified:*

15: sp_rvirus:*

16: sp_bacteriap:*

17: sp_archeap:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
No.	Score	Match	Length	DB	ID	
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1	3883	83.0	855	11	Q9JJI7	Q9jji7 rattus norv
2	2664	56.9	845	13	Q9DGR1	Q9dgr1 xenopus lae
3	2379	50.8	422	4	Q8WVC1	Q8wvc1 homo sapien
4	1011.5	21.6	572	11	Q8BIK6	Q8bik6 mus musculu
5	717.5	15.3	855	4	Q7Z410	Q7z410 homo sapien
6	717.5	15.3	1059	4	Q7Z411	Q7z411 homo sapien
7	690.5	14.8	1111	11	Q80YN4	Q80yn4 rattus norv
8	687	14.7	777	11	Q8CAN9	Q8can9 mus musculu
9	644.5	13.8	767	13	Q9DGR2	Q9dgr2 xenopus lae
10	636.5	13.6	680	5	Q868H7	Q868h7 branchiosto
11	623.5	13.3	680	5	Q868H5	Q868h5 branchiosto
12	617	13.2	581	5	Q9XZM7	Q9xzm7 strongyloce
13	612	13.1	688	5	Q868H6	Q868h6 branchiosto
14	601	12.8	490	11	Q7TN04	Q7tn04 mus musculu
15	600	12.8	490	11	Q920K3	Q920k3 rattus norv

Database : Published_Applications_AA:*

1: /cgn2_6/ptodata/2/pubpaa/US07_PUBCOMB.pep:*

2: /cgn2_6/ptodata/2/pubpaa/PCT_NEW_PUB.pep:*

3: /cgn2_6/ptodata/2/pubpaa/US06_NEW_PUB.pep:*

4: /cgn2_6/ptodata/2/pubpaa/US06_PUBCOMB.pep:*

5: /cgn2_6/ptodata/2/pubpaa/US07_NEW_PUB.pep:*

6: /cgn2_6/ptodata/2/pubpaa/PCTUS_PUBCOMB.pep:*

7: /cgn2_6/ptodata/2/pubpaa/US08_NEW_PUB.pep:*

8: /cgn2_6/ptodata/2/pubpaa/US08_PUBCOMB.pep:*

9: /cgn2_6/ptodata/2/pubpaa/US09A_PUBCOMB.pep:*

10: /cgn2_6/ptodata/2/pubpaa/US09B_PUBCOMB.pep:*

11: /cgn2_6/ptodata/2/pubpaa/US09C_PUBCOMB.pep:*

12: /cgn2_6/ptodata/2/pubpaa/US09_NEW_PUB.pep:*

13: /cgn2_6/ptodata/2/pubpaa/US10A_PUBCOMB.pep:*

14: /cgn2_6/ptodata/2/pubpaa/US10B_PUBCOMB.pep:*

15: /cgn2_6/ptodata/2/pubpaa/US10C_PUBCOMB.pep:*

16: /cgn2_6/ptodata/2/pubpaa/US10_NEW_PUB.pep:*

17: /cgn2_6/ptodata/2/pubpaa/US60_NEW_PUB.pep:*

18: /cgn2_6/ptodata/2/pubpaa/US60_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
	No.	Score	Match	Length	DB ID	
1	4681	100.0	855	10	US-09-776-191-2	Sequence 2, Appli
2	4681	100.0	855	12	US-10-072-012-352	Sequence 352, App
3	4681	100.0	855	12	US-10-072-012-411	Sequence 411, App
4	4681	100.0	855	12	US-10-072-012-418	Sequence 418, App
5	4681	100.0	855	14	US-10-099-700A-2	Sequence 2, Appli
6	4681	100.0	855	14	US-10-190-030B-2	Sequence 2, Appli
7	4681	100.0	855	14	US-10-302-840A-2	Sequence 2, Appli
8	4681	100.0	855	14	US-10-267-219-2	Sequence 2, Appli
9	4681	100.0	855	14	US-10-112-221A-2	Sequence 2, Appli
10	4681	100.0	855	14	US-10-104-271-2	Sequence 2, Appli
11	4681	100.0	855	15	US-10-147-211A-2	Sequence 2, Appli
12	4681	100.0	855	15	US-10-156-214A-2	Sequence 2, Appli
13	4681	100.0	855	16	US-10-600-187-2	Sequence 2, Appli
14	4676	99.9	855	12	US-10-072-012-353	Sequence 353, App
15	4676	99.9	855	12	US-10-072-012-412	Sequence 412, App
16	4676	99.9	855	12	US-10-072-012-419	Sequence 419, App
17	4676	99.9	855	15	US-10-295-027-1185	Sequence 1185, Ap
18	4672	99.8	855	12	US-10-072-012-354	Sequence 354, App
19	4672	99.8	855	12	US-10-072-012-420	Sequence 420, App
20	4672	99.8	855	12	US-10-037-417-132	Sequence 132, App
21	4631	98.9	851	12	US-10-276-774-1798	Sequence 1798, Ap
22	4631	98.9	851	12	US-10-296-115-1143	Sequence 1143, Ap
23	4175.5	89.2	782	14	US-10-097-340-312	Sequence 312, App
24	4175	89.2	762	16	US-10-729-807-1	Sequence 1, Appli
25	4111	87.8	757	12	US-10-072-012-44	Sequence 44, Appli
26	3901	83.3	855	9	US-09-900-751-2	Sequence 2, Appli
27	3901	83.3	855	12	US-10-072-012-355	Sequence 355, App
28	3901	83.3	855	12	US-10-072-012-413	Sequence 413, App

29	3883	83.0	855	12	US-10-072-012-356	Sequence 356, App
30	3883	83.0	855	12	US-10-072-012-414	Sequence 414, App
31	3883	83.0	855	12	US-10-072-012-417	Sequence 417, App
32	3810	81.4	902	12	US-10-333-743-3	Sequence 3, Appli
33	3810	81.4	902	16	US-10-600-187-10	Sequence 10, Appl
34	3810	81.4	902	16	US-10-297-987B-11	Sequence 11, Appl
35	2980	63.7	620	9	US-09-925-301-1193	Sequence 1193, Ap
36	2664	56.9	845	12	US-10-072-012-415	Sequence 415, App
37	1319	28.2	241	10	US-09-776-191-50	Sequence 50, Appl
38	1319	28.2	241	14	US-10-099-700A-4	Sequence

Database : SwissProt_42:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
No.	Score	Match	Length	DB	ID	
1	4676	99.9	855	1	ST14_HUMAN	Q9y5y6 homo sapien
2	3901	83.3	855	1	ST14_MOUSE	P56677 mus musculu
3	1124.5	24.0	811	1	TMS6_MOUSE	Q9dbi0 mus musculu
4	1124	24.0	811	1	TMS6_HUMAN	Q8iu80 homo sapien
5	727	15.5	1034	1	ENTK_PIG	P98074 sus scrofa
6	712.5	15.2	1035	1	ENTK_BOVIN	P98072 bos taurus
7	692	14.8	1042	1	CORI_HUMAN	Q9y5q5 homo sapien
8	682.5	14.6	1019	1	ENTK_HUMAN	P98073 homo sapien
9	676.5	14.5	1069	1	ENTK_MOUSE	P97435 mus musculu
10	663.5	14.2	1113	1	CORI_MOUSE	Q9z319 mus musculu
11	600	12.8	490	1	TMS2_MOUSE	Q9jiq8 mus musculu
12	588	12.6	422	1	DES1_HUMAN	Q9ul52 homo sapien
13	586.5	12.5	704	1	CRAR_MOUSE	P98064 mus musculu
14	574	12.3	699	1	CRAR_HUMAN	P48740 h complemen
15	558.5	11.9	492	1	TMS2_HUMAN	O15393 homo sapien
16	546	11.7	453	1	TMS3_MOUSE	Q8k1t0 mus musculu
17	533.5	11.4	638	1	KAL_MOUSE	P26262 mus musculu
18	533	11.4	454	1	TMS3_HUMAN	P57727 homo sapien
19	518	11.1	603	1	CFAI_MOUSE	Q61129 mus musculu
20	518	11.1	604	1	CFAI_RAT	Q9wuw3 rattus norv
21	514.5	11.0	638	1	KAL_RAT	P14272 rattus norv
22	513	11.0	455	1	TMS5_MOUSE	Q9er04 mus musculu
23	511.5	10.9	418	1	HATT_HUMAN	O60235 homo sapien

Database : A_Geneseq_29Jan04:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:*

6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
No.	Score	Match	Length	DB	ID	
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1	4681	100.0	855	2	AAY06671	Aay06671 Tumour an
2	4681	100.0	855	4	AAB98500	Aab98500 Human TAD
3	4681	100.0	855	4	AAE06930	Aae06930 Human mem
4	4681	100.0	855	5	AAO22929	Aao22929 Type II t
5	4681	100.0	855	6	ABP56619	Abp56619 Human mem
6	4681	100.0	855	6	AAO30146	Aao30146 Human mem
7	4681	100.0	855	6	AAE29820	Aae29820 Human mem
8	4681	100.0	855	6	AAE29791	Aae29791 Human mem
9	4681	100.0	855	6	ABP72376	Abp72376 Transmemb
10	4681	100.0	855	7	ADB97551	Adb97551 Human MTS
11	4676	99.9	855	3	AAB19552	Aab19552 Human mat
12	4676	99.9	855	4	AAB35465	Aab35465 Human mem
13	4631	98.9	851	4	AAM25628	Aam25628 Human pro
14	4631	98.9	851	4	ABB11428	Abb11428 Human mem
15	4319	92.3	932	4	ABG21442	Abg21442 Novel hum
16	4175.5	89.2	782	5	ABG96427	Abg96427 Human ova
17	4175	89.2	762	3	AAY90284	Aay90284 Human pep
18	3901	83.3	855	5	AAE23083	Aae23083 Epithin p
19	3810	81.4	902	4	AAB98507	Aab98507 Murine ep
20	3810	81.4	902	5	AAU80517	Aau80517 Mouse epi
21	3810	81.4	902	5	AAU77549	Aau77549 Murine ty
22	3781	80.8	683	3	AAB19551	Aab19551 Human mat
23	2980	63.7	620	3	AAB43748	Aab43748 Human can
24	1352	28.9	362	4	ABG21441	Abg21441 Novel hum

Database : PIR_78:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result	Query					Description
No.	Score	Match	Length	DB	ID	
1	3883	83.0	855	2	JC7731	membrane-bound arg
2	727	15.5	1034	1	A53663	enteropeptidase (E
3	712.5	15.2	1035	1	A43090	enteropeptidase (E
4	682.5	14.6	1019	1	A56318	enteropeptidase (E
5	663.5	14.2	1113	2	JE0315	low-density lipopr
6	578.5	12.4	1524	2	T30337	polyprotein - Afri
7	574	12.3	699	1	I54763	Ra-reactive factor
8	533.5	11.4	638	1	KQMSPL	plasma kallikrein
9	514.5	11.0	638	1	KQRTPL	plasma kallikrein
10	509.5	10.9	790	1	PLPG	plasmin (EC 3.4.21
11	506	10.8	613	2	S15468	complement C3b/C4b
12	502	10.7	460	2	B61545	plasmin (EC 3.4.21
13	501.5	10.7	786	1	A47547	serine proteinease
14	500	10.7	638	1	KQHUP	plasma kallikrein
15	497	10.6	810	1	PLHU	plasmin (EC 3.4.21
16	492.5	10.5	583	2	A29154	complement factor
17	491.5	10.5	812	1	PLMS	plasmin (EC 3.4.21
18	491	10.5	416	1	KFBO	coagulation factor
19	490.5	10.5	812	1	PLBO	plasmin (EC 3.4.21
20	490	10.5	417	1	S00845	hepsin (EC 3.4.21

RESULT 1

Q9JJI7

ID Q9JJI7 PRELIMINARY; PRT; 855 AA.
AC Q9JJI7;
DT 01-OCT-2000 (TrEMBLrel. 15, Created)
DT 01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
DT 01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE Membrane bound serine protease (Membrane bound arginine specific
DE serine protease).
GN MBSP.
OS Rattus norvegicus (Rat).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX NCBI_TaxID=10116;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=WISTAR; TISSUE=Jejunum;
RA Tsuzuki S.;
RT "A membrane bound serine protease expressed in rat small intestine.";
RL Submitted (JAN-2000) to the EMBL/GenBank/DDBJ databases.
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN=Wistar; TISSUE=Duodenum;
RA Inoue H., Takahashi K., Kishi K.;
RT "membrane-bound arginine specific serine protease.";
RL Submitted (SEP-2000) to the EMBL/GenBank/DDBJ databases.
CC -!- SIMILARITY: BELONGS TO PEPTIDASE FAMILY S1.
CC -!- SIMILARITY: CONTAINS 2 CUB DOMAINS.
DR EMBL; AB037898; BAB03502.1; -.
DR EMBL; AB049189; BAB13765.1; -.
DR PIR; JC7731; JC7731.
DR HSSP; P00763; 1DPO.
DR MEROPS; S01.302; -.
DR GO; GO:0004263; F:chymotrypsin activity; IEA.
DR GO; GO:0008233; F:peptidase activity; IEA.
DR GO; GO:0004295; F:trypsin activity; IEA.
DR GO; GO:0006508; P:proteolysis and peptidolysis; IEA.
DR InterPro; IPR000859; CUB.
DR InterPro; IPR009003; Cys_Ser_trypsin.
DR InterPro; IPR002172; LDL_receptor_A.
DR InterPro; IPR001254; Peptidase_S1.
DR InterPro; IPR001314; Peptidase_S1A.
DR Pfam; PF00431; CUB; 2.
DR Pfam; PF00057; ldl_recept_a; 4.
DR Pfam; PF00089; trypsin; 1.
DR PRINTS; PR00722; CHYMOTRYPSIN.
DR PRINTS; PR00261; LDLRECEPTOR.
DR SMART; SM00042; CUB; 2.
DR SMART; SM00192; LDLa; 3.
DR SMART; SM00020; Tryp_SPC; 1.
DR PROSITE; PS01180; CUB; 2.
DR PROSITE; PS01209; LDLRA_1; 2.
DR PROSITE; PS50068; LDLRA_2; 4.
DR PROSITE; PS50240; TRYPSIN_DOM; 1.
DR PROSITE; PS00134; TRYPSIN_HIS; 1.
DR PROSITE; PS00135; TRYPSIN_SER; 1.
KW Hydrolase; Protease; Serine protease.

Db	721	AEYSTVVRPICLPDNTHVFPAGKAIWVTGWGHTKEGGTGALILQKGEIRVINQTTCEELL	780
Qy	781	PQQITPRMMCVCVGFLSGGVVDSCQGDSGGPLSSVEADGRIFQAGVVSWGDGCAQRNKGPGVYT	840
		:	
Db	781	PQQITPRMMCVCVGFLSGGVVDSCQGDSGGPLSSVEKDGRIFQAGVVSWGEKGCAQRNKGPGVYT	840
Qy	841	RLPLFRDWIKENTGV	855
		:	
Db	841	RIPEVRDWIKEQTGV	855

RESULT 2

ST14_MOUSE

ID ST14_MOUSE STANDARD; PRT; 855 AA.
AC P56677;
DT 15-JUL-1999 (Rel. 38, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Suppressor of tumorigenicity 14 (EC 3.4.21.-) (Epithin).
GN ST14 OR PRSS14.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C.B.17SCID; TISSUE=Thymus;
RX MEDLINE=99216440; PubMed=10199918;
RA Kim M.G., Chen C., Lyu M.S., Cho E.G., Park D., Kozak C.,
RA Schwartz R.H.;
RT "Cloning and chromosomal mapping of a gene isolated from thymic
RT stromal cells encoding a new mouse type II membrane serine protease,
RT epithin, containing four LDL receptor modules and two CUB domains.";
RL Immunogenetics 49:420-428(1999).
RN [2]
RP REVISIONS TO 23; 321; 325; 343; 409-410 AND C-TERMINUS.
RC STRAIN=C.B.17SCID; TISSUE=Thymus;
RA Kim M.G., Chen C., Cho E.G., Park D., Schwartz R.H.;
RL Submitted (MAR-2000) to the EMBL/GenBank/DDBJ databases.
RN [3]
RP SEQUENCE FROM N.A.
RC TISSUE=Breast tumor;
RX MEDLINE=22388257; PubMed=12477932;
RA Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
RA Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RA Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
RA Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
RA Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
RA Schnurch A., Schein J.E., Jones S.J.M., Marra M.A.;
RT "Generation and initial analysis of more than 15,000 full-length
RT human and mouse cDNA sequences.";
RL Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
CC -!- SUBCELLULAR LOCATION: Type II membrane protein (Probable).
CC -!- TISSUE SPECIFICITY: Highly expressed in intestine, kidney, lung,
CC and thymus. Not expressed in skeletal muscle, liver, heart,
CC testis and brain.
CC -!- SIMILARITY: Belongs to peptidase family S1.

CC -!- SIMILARITY: Contains 2 CUB domains.
CC -!- SIMILARITY: Contains 4 LDL-receptor class A domains.
CC -----
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
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CC -----
DR EMBL; AF042822; AAD02230.3; -.
DR EMBL; BC005496; AAH05496.1; -.
DR HSSP; P20231; 1AAO.
DR MEROPS; S01.302; -.
DR MGD; MGI:1338881; St14.
DR GO; GO:0005576; C:extracellular; IDA.
DR GO; GO:0019897; C:extrinsic to plasma membrane; IDA.
DR GO; GO:0008236; F:serine-type peptidase activity; IDA.
DR InterPro; IPR000859; CUB.
DR InterPro; IPR009003; Cys_Ser_trypsin.
DR InterPro; IPR002172; LDL_receptor_A.
DR InterPro; IPR001254; Peptidase_S1.
DR InterPro; IPR001314; Peptidase_S1A.
DR Pfam; PF00431; CUB; 2.
DR Pfam; PF00057; ldl_recept_a; 4.
DR Pfam; PF00089; trypsin; 1.
DR PRINTS; PR00722; CHYMOTRYPSIN.
DR PRINTS; PR00261; LDLRECEPTOR.
DR SMART; SM00042; CUB; 2.
DR SMART; SM00192; LDLa; 4.
DR SMART; SM00020; Tryp_SPC; 1.
DR PROSITE; PS01180; CUB; 2.
DR PROSITE; PS01209; LDLRA_1; 2.
DR PROSITE; PS50068; LDLRA_2; 4.
DR PROSITE; PS50240; TRYPSIN_DOM; 1.
DR PROSITE; PS00134; TRYPSIN_HIS; 1.
DR PROSITE; PS00135; TRYPSIN_SER; 1.
KW Signal-anchor; Glycoprotein; Hydrolase; Serine protease;
KW Transmembrane; Repeat.
FT DOMAIN 1 55 CYTOPLASMIC (POTENTIAL).
FT TRANSMEM 56 76 SIGNAL-ANCHOR (TYPE-II MEMBRANE PROTEIN)
(POTENTIAL).
FT DOMAIN 77 855 EXTRACELLULAR (POTENTIAL).
FT DOMAIN 214 331 CUB 1.
FT DOMAIN 340 444 CUB 2.
FT DOMAIN 451 488 LDL-RECEPTOR CLASS A 1.
FT DOMAIN 489 522 LDL-RECEPTOR CLASS A 2.
FT DOMAIN 523 561 LDL-RECEPTOR CLASS A 3.
FT DOMAIN 565 604 LDL-RECEPTOR CLASS A 4.
FT DOMAIN 615 854 SERINE PROTEASE.
FT ACT_SITE 656 656 CHARGE RELAY SYSTEM (BY SIMILARITY).
FT ACT_SITE 711 711 CHARGE RELAY SYSTEM (BY SIMILARITY).
FT ACT_SITE 805 805 CHARGE RELAY SYSTEM (BY SIMILARITY).
FT CARBOHYD 107 107 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 302 302 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 365 365 N-LINKED (GLCNAC. . .) (POTENTIAL).

FT CARBOHYD 421 421 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CARBOHYD 489 489 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CARBOHYD 772 772 N-LINKED (GLCNAC. . .) (POTENTIAL).
 SQ SEQUENCE 855 AA; 94654 MW; 4F10E84DA2146DD5 CRC64;

 Query Match 83.3%; Score 3901; DB 1; Length 855;
 Best Local Similarity 81.8%; Pred. No. 6.2e-261;
 Matches 699; Conservative 73; Mismatches 83; Indels 0; Gaps 0;

 Qy 1 MGSDRARKGGGGPKDFGAGLKYNRSRHEKVNGLEEGVEFLPVNNVKKVEKGPGRWVVLAA 60
 ||||:||| ||| :||||||| ||| :||| ||||| ||| ||||| ||| ||||| ||| ||||| |||
 Db 1 MGSNRGRKAGGGSQDFGAGLKYNRSLENMNGFEEGVEFLPANNAKKVEKRGPRRWVVLVA 60

 Qy 61 VLIGLLLVLIGFLVWHLQYRDVRVQKVFNNGYMRITNENFDAYENSNSTEFVSLASKV 120
 ||| |||:| | | ||| |||:||||||| |||:| | | ||| |||:| | | |||:| | | :| |
 Db 61 VLFSFLLLSLMAGLLVWHFHYRNVRVQKVFNNGHLRITNEIFLDAYENSTSTEFISLASQV 120

 Qy 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYYWSEFSIPQHLVEEAERVMAERVVM 180
 ||:|||||:||| |||:|||||||:||||||| |||:||||||| |||:| | | ||| :| | | |||
 Db 121 KEALKLLYNEVPVLGPYHKESAVTAFSEGSVIAYYWSEFSIPPHLAEEVDRAMAVERVVT 180

 Qy 181 LPPRARSLKSFVVTSVVAFPTDSKTVQRTQDNCSCFGLHARGVELMRFTTPGFPDSPYPA 240
 ||||||:|||||:||||||| ||| :| :||||||| ||| ||| :| |||:|||||:|||||||
 Db 181 LPPRARALKSFVLTSVVAFPIDPRMLQRTQDNCSCFALHAHGAATRFTTPGFPNSPYPA 240

 Qy 241 HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYNTLSPMEHALVQLCGTYPPS 300
 ||||||| ||||||| ||||||| |||:| ||| |||:|||||||:| |||:|||||||:|:|||||:|||
 Db 241 HARCQWVLRGDADSVLSLTFRSFDVAPCDEHGSDLVTYDLSLSPMEPHAVVRLCGTFSPS 300

 Qy 301 YNLTFHSSQNVLLITLITNTERRHPGFEATFFQLPRMSSCGGRLRKAQGTFNSPYYPGHY 360
 ||||||| |||:| |||:| |||:| |||:| |||:| |||:| |||:| |||:| |||:| |||:|
 Db 301 YNLTFLSSQNVFLVTLITNTDRRHPGFEATFFQLPKMSSCGGFLSDTQGTFSSPYYPGHY 360

 Qy 361 PPNIDCTWNIEVPNNQHVKVSKFFYLLEPGVPAGTCPKDVEINGEKGERSQFVVTS 420
 ||||:|||||:|||||:| ||| |||:| |||:| |||:| |||:| |||:| |||:| |||:|
 Db 361 PPNINCTWNIKVPNNRNVKVRFKLFYLVDPNVPVGSTKDYVEINGEKGERSQFVVSS 420

 Qy 421 NSNKITVRFHSDQSYTDTGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCDGWADCTDH 480
 |||:||||| ||| ||| ||| ||| ||| |||:| ||| |||:| |||:| |||:| |||:|
 Db 421 NSSKITVHFHSDHSYTDTGFLAEYLSYDSNDPCPGFMCKTGRCTIRKELRCDGWADCPDY 480

 Qy 481 SDELNCSCDAGHQFTCKNFKPLFWVCDSVNDGDSDEQGCSCPAQTFRCNSNGKCLSK 540
 ||| | | :| ||| |||:| ||| ||| ||| ||| ||| |||:| |||:| |||:| :| |||:|
 Db 481 SDERYCRCNATHQFTCKNQFCKPLFWVCDSVNDGDSDEEGCSCPAGSFKCSNGKCLPQ 540

 Qy 541 SQQCNGKDDCGDSDEASCPKVNVVTCTKHTYRCLNGLCLSKGNPECDGKEDCSDGSDEK 600
 |||:|||||:||||||| |||:| |||:| |||:| |||:| |||:| |||:| |||:|
 Db 541 SQKCNGKDNCGDSDEASCDSVNVVSCTKYTYRCQNGLCLSKGNPECDGKTDSDGSDEK 600

 Qy 601 DCDCGLRSFTRQARVVGTDADEGEWPWQVSLHALGQGHICGASLISPWLVSAAHCYID 660
 :| ||| |||:| ||| |||:| ||| ||| ||| |||:| ||| |||:| ||| |||:|
 Db 601 NDCGGLRSFTKQARVVGTTADEGEWPWQVSLHALGQGHLCGASLISPWLVSAAHCFQD 660

 Qy 661 DRGFRYSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLELEKP 720
 |:| :| | | ||| ||| |||:| |||:| |||:| |||:| |||:| |||:|
 Db 661 DKNFKYSDYTMWTAFLGLLDQSKRSASGVQELKLKRIITHPSFNDFTFDYDIALLELEKS 720

Qy 721 AEYSSMVRPICLPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCEPLL 780
|||:|||||||:|||||||:|||||||:|||||||:|||||||:|||||||:|||:
Db 721 VEYSTVVRPICLPDATHVFPAGKAIWVTGWGHTKEGGTGALILQKGEIRVINQTTCEDLM 780

Qy 781 PQQITPRMMCVGFLSGGVVDSCQGDSGGPLSSVEADGRIFQAGVVSWGDGCAQRNKGPGVYT 840
|||||:|||||||:|||||||:|||||||:|||||||:|||||||:
Db 781 PQQITPRMMCVGFLSGGVVDSCQGDSGGPLSSAEKDGRMFQAGVVSWGEGCAQRNKGPGVYT 840

Qy 841 RLPLFRDWIKENTGV 855
|||:|||||||:|||:
Db 841 RLPVVVRDWISEHTGV 855

RESULT 1

ST14_HUMAN

ID ST14_HUMAN STANDARD; PRT; 855 AA.
AC Q9Y5Y6; Q9BS01; Q9H3S0; Q9HB36; Q9HCA3;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Suppressor of tumorigenicity 14 (EC 3.4.21.-) (Matriptase) (Membrane-
DE type serine protease 1) (MT-SP1) (Prostamin) (Serine protease TADG-15)
DE (Tumor associated differentially-expressed gene-15 protein).
GN ST14 OR PRSS14 OR SNC19 OR TADG15.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99303581; PubMed=10373424;
RA Lin C.Y., Anders J., Johnson M., Sang Q.A., Dickson R.B.;
RT "Molecular cloning of cDNA for matriptase, a matrix-degrading serine
RT protease with trypsin-like activity.";
RL J. Biol. Chem. 274:18231-18236 (1999).
RN [2]
RP SEQUENCE FROM N.A.
RX MEDLINE=99432178; PubMed=10500122;
RA Takeuchi T., Shuman M.A., Craik C.S.;
RT "Reverse biochemistry: Use of macromolecular protease inhibitors to
RT dissect complex biological processes and identify a membrane-type
RT serine protease in epithelial cancer and normal tissue.";
RL Proc. Natl. Acad. Sci. U.S.A. 96:11054-11061 (1999).
RN [3]
RP SEQUENCE FROM N.A.
RC TISSUE=Prostate;
RA Yamaguchi N., Mitsui S.;
RT "Molecular cloning of a novel transmembrane serine protease expressed
RT in human prostate.";
RL Submitted (JUL-1999) to the EMBL/GenBank/DDBJ databases.
RN [4]
RP SEQUENCE FROM N.A.
RA Tanimoto H., Underwood L.J., Wang Y., Shigemasa K., Parmley T.H.,
RA O'Brien T.J.;
RL Submitted (APR-1998) to the EMBL/GenBank/DDBJ databases.
RN [5]
RP SEQUENCE FROM N.A.
RC TISSUE=Blood, and Muscle;
RX MEDLINE=22388257; PubMed=12477932;
RA Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
RA Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RA Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,

RA Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
RA Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
RA Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RT "Generation and initial analysis of more than 15,000 full-length
RT human and mouse cDNA sequences.";
RL Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903 (2002).
RN [6]
RP SEQUENCE OF 340-664 FROM N.A.
RA Cao J., Fan W., Zheng S.;
RT "Genomic analysis of a novel human serine protease SNC19.";
RL Submitted (JUN-2000) to the EMBL/GenBank/DDBJ databases.
RN [7]
RP CHARACTERIZATION.
RC TISSUE=Milk;
RX MEDLINE=99303582; PubMed=10373425;
RA Lin C.Y., Anders J., Johnson M., Dickson R.B.;
RT "Purification and characterization of a complex containing matriptase
RT and a Kunitz-type serine protease inhibitor from human milk.";
RL J. Biol. Chem. 274:18237-18242 (1999).
CC -!- FUNCTION: Degrades extracellular matrix. Proposed to play a role
CC in breast cancer invasion and metastasis. Exhibits trypsin-like
CC activity as defined by cleavage of synthetic substrates with Arg
CC or Lys as the P1 site.
CC -!- SUBCELLULAR LOCATION: Type II membrane protein (Probable).
CC -!- SIMILARITY: Belongs to peptidase family S1.
CC -!- SIMILARITY: Contains 2 CUB domains.
CC -!- SIMILARITY: Contains 4 LDL-receptor class A domains.
CC -----
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC the European Bioinformatics Institute. There are no restrictions on its
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CC or send an email to license@isb-sib.ch).
CC -----
DR EMBL; AF118224; AAD42765.2; -.
DR EMBL; AF133086; AAF00109.1; -.
DR EMBL; AB030036; BAB20376.1; -.
DR EMBL; AF057145; AAG15395.1; -.
DR EMBL; BC005826; AAH05826.1; -.
DR EMBL; BC030532; AAH30532.1; -.
DR EMBL; AF283256; AAG13949.1; -.
DR HSSP; P00763; 1DPO.
DR Genew; HGNC:11344; ST14.
DR MIM; 606797; -.
DR MEROPS; S01.302; -.
DR GO; GO:0005887; C:integral to plasma membrane; TAS.
DR GO; GO:0008236; F:serine-type peptidase activity; TAS.
DR GO; GO:0006508; P:proteolysis and peptidolysis; TAS.
DR InterPro; IPR000859; CUB.
DR InterPro; IPR009003; Cys_Ser_trypsin.
DR InterPro; IPR002172; LDL_receptor_A.
DR InterPro; IPR001254; Peptidase_S1.

DR InterPro; IPR001314; Peptidase_S1A.
 DR Pfam; PF00431; CUB; 2.
 DR Pfam; PF00057; ldl_recept_a; 4.
 DR Pfam; PF00089; trypsin; 1.
 DR PRINTS; PR00722; CHYMOTRYPSIN.
 DR PRINTS; PR00261; LDLRECEPTOR.
 DR SMART; SM00042; CUB; 2.
 DR SMART; SM00192; LDLa; 3.
 DR SMART; SM00020; Tryp_SPC; 1.
 DR PROSITE; PS01180; CUB; 2.
 DR PROSITE; PS01209; LDLRA_1; 2.
 DR PROSITE; PS50068; LDLRA_2; 4.
 DR PROSITE; PS50240; TRYPSIN_DOM; 1.
 DR PROSITE; PS00134; TRYPSIN_HIS; 1.
 DR PROSITE; PS00135; TRYPSIN_SER; 1.
 KW Signal-anchor; Glycoprotein; Hydrolase; Serine protease;
 KW Transmembrane; Repeat.
 FT DOMAIN 1 55 CYTOPLASMIC (POTENTIAL).
 FT TRANSMEM 56 76 SIGNAL-ANCHOR (TYPE-II MEMBRANE PROTEIN)
 (POTENTIAL).
 FT DOMAIN 77 855 EXTRACELLULAR (POTENTIAL).
 FT DOMAIN 214 334 CUB 1.
 FT DOMAIN 340 447 CUB 2.
 FT DOMAIN 452 487 LDL-RECEPTOR CLASS A 1.
 FT DOMAIN 487 524 LDL-RECEPTOR CLASS A 2.
 FT DOMAIN 524 560 LDL-RECEPTOR CLASS A 3.
 FT DOMAIN 566 603 LDL-RECEPTOR CLASS A 4.
 FT DOMAIN 615 854 SERINE PROTEASE.
 FT ACT_SITE 656 656 CHARGE RELAY SYSTEM (BY SIMILARITY).
 FT ACT_SITE 711 711 CHARGE RELAY SYSTEM (BY SIMILARITY).
 FT ACT_SITE 805 805 CHARGE RELAY SYSTEM (BY SIMILARITY).
 FT CARBOHYD 109 109 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CARBOHYD 302 302 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CARBOHYD 485 485 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CARBOHYD 772 772 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT CONFLICT 327 329 FEA -> GTR (IN REF. 5; AAH05826).
 FT CONFLICT 381 381 R -> S (IN REF. 4).
 FT CONFLICT 674 674 A -> V (IN REF. 3).
 SQ SEQUENCE 855 AA; 94769 MW; 26143132C01F99C9 CRC64;

Query Match 99.9%; Score 4676; DB 1; Length 855;
 Best Local Similarity 99.9%; Pred. No. 2.7e-314;
 Matches 854; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

Qy 1 MGSDRARKGGGGPKDFGAGLKYNNSRHEKVNGLEEGVEFLPVNNVKVKEHGPGRWVVLAA 60
 |||||||
 Db 1 MGSDRARKGGGGPKDFGAGLKYNNSRHEKVNGLEEGVEFLPVNNVKVKEHGPGRWVVLAA 60

Qy 61 VLIGLLLVLIGFLVWHLQYRDVRVQKVFNGYMRITNENFDAYENSNSTEFVSLASKV 120
 |||||||
 Db 61 VLIGLLLVLIGFLVWHLQYRDVRVQKVFNGYMRITNENFDAYENSNSTEFVSLASKV 120

Qy 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM 180
 |||||||
 Db 121 KDALKLLYSGVPFLGPYHKESAVTAFSEGSVIAYWSEFSIPQHLVEEAERVMAERVVM 180

Qy 181 LPPRARSLKSFVVTSVVAFPTDSKTVQRTQDNSCSFGLHARGVELMRFTPGFPDSPYPA 240

Db ||||||| 181 LPPRARSLKFVVTSVVAFPTDSKTVQRTQDNCSFGLHARGVELMRFTTPGFPDSPYPA 240

Qy ||||||| 241 HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYVNTLSPMEPHALVQLCGTYPPS 300

Db ||||||| 241 HARCQWALRGDADSVLSLTFRSFDLASCDERGSDLVTYVNTLSPMEPHALVQLCGTYPPS 300

Qy ||||||| 301 YNLTFHSSQNVLLITLITNTERRHPFEATFFQLPRMSSCGGRLRKAQGTFNSPYPYGHY 360

Db ||||||| 301 YNLTFHSSQNVLLITLITNTERRHPFEATFFQLPRMSSCGGRLRKAQGTFNSPYPYGHY 360

Qy ||||||| 361 PPNIDCTWNIEVPNNQHVVKVSFKFFYLLEPGVPAGTCPKDVEINGEKYCGERSQFVVTS 420

Db ||||||| 361 PPNIDCTWNIEVPNNQHVVKVRFKFFYLLEPGVPAGTCPKDVEINGEKYCGERSQFVVTS 420

Qy ||||||| 421 NSNKITVRFHSDQSYTDTGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCGWADCTDH 480

Db ||||||| 421 NSNKITVRFHSDQSYTDTGFLAEYLSYDSSDPCPGQFTCRTGRCIRKELRCGWADCTDH 480

Qy ||||||| 481 SDELNCSCDAGHQFTCKNKFKPLFWVCDSVNDGDNSEQGCSCPAQTFRCNSNGKCLSK 540

Db ||||||| 481 SDELNCSCDAGHQFTCKNKFKPLFWVCDSVNDGDNSEQGCSCPAQTFRCNSNGKCLSK 540

Qy ||||||| 541 SQQCNGKDDCGDGSEASCPKVNVTCTKHTYRCLNGLCLSKGNECDGKEDCSDGSDEK 600

Db ||||||| 541 SQQCNGKDDCGDGSEASCPKVNVTCTKHTYRCLNGLCLSKGNECDGKEDCSDGSDEK 600

Qy ||| 601 DCDCGLRSFTRQARVVGTDADEGEWPWQVSLHALGQGHICGASLISPWLVSAAHCYID 660

Db ||| 601 DCDCGLRSFTRQARVVGTDADEGEWPWQVSLHALGQGHICGASLISPWLVSAAHCYID 660

Qy ||| 661 DRGFYSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP 720

Db ||| 661 DRGFYSDPTQWTAFLGLHDQSQRSAAPGVQERRLKRIISHPFFNDFTFDYDIALLEKP 720

Qy ||| 721 AEYSSMVRPICLDPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCE 780

Db ||| 721 AEYSSMVRPICLDPDASHVFPAGKAIWVTGWGHTQYGGTGALILQKGEIRVINQTTCE 780

Qy ||| 781 PQQITPRMMCVGFLSGGVVDSCQGD¹⁹⁰SGGPLSSVEADGRIFQAGVVSWGDGCAQRNKG²⁰⁰PGVYT 840

Db ||| 781 PQQITPRMMCVGFLSGGVVDSCQGD¹⁹⁰SGGPLSSVEADGRIFQAGVVSWGDGCAQRNKG²⁰⁰PGVYT 840

Qy ||| 841 RLPLFRDWIKENTGV 855

Db ||| 841 RLPLFRDWIKENTGV 855